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AMENDMENTS TO THE CLAIMS

This listing of claims will replace all prior versions, and listings, of the claims in the application:

Listing of Claims:

1. (Currently amended) A method for amplifying expressed genetic sequences from genomic DNA (gDNA) selected from a higher-order eukaryotic species, for printing on DNA microarrays, wherein the method comprises:

identifying a 3' untranslated region (3'UTR) of a gDNA sequence based on the presence of a stop codon and a polyadenylation signal in the gDNA sequence;

selecting a predetermined gDNA sequence within the 3'UTR;

designing a probe for said predetermined gDNA sequence;

performing a first polymerase chain reaction (PCR) for the 3'UTR on gDNA to generate a first PCR-product;

separating the resultant-resulting first PCR-product by a size-differentiation process selected from the group consisting of electrophoresis and chromatography;

selecting a predetermined band from the size-differentiated samples;

performing a second polymerase chain reaction to amplify a PCR product in the predetermined band; and the selected predetermined band to generate a second PCR product; and

depositing a sequence amplified by said second polymerase chain reaction to a substrate of an array.

printing the second PCR product on a substrate to form an array, wherein the printed product is free of polyadenosine sequences.

2. - 3. (Cancel)

4. (Previously presented) The method according to claim 1, wherein said predetermined gDNA sequence within the 3'UTR is selected by use of computer software.

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- 5. (Previously presented) The method according to claim 1, wherein said selected predetermined gDNA sequence within the 3'UTR has a length of at least about 75 nucleotides.
- 6. (Original) The method according to claim 5, wherein said selected predetermined gDNA sequence has a length of about 200 to about 600 bases.
- 7. (Original) The method according to claim 6, wherein said selected predetermined gDNA sequence has a length of about 250 to about 450 bases.
- 8. (Original) The method according to claim 1, wherein said selected predetermined gDNA sequence has an overall homology of less than or equal to about 70% to any other genomic sequence in the same genome.
- 9. (Original) The method according to claim 8, wherein said selected predetermined gDNA sequence has an overall homology of less than or equal to about 40% to any other genomic sequence in the same genome.
- 10. (Original) The method according to claim 8, wherein said selected predetermined gDNA sequence has an overall homology of from about 20% to 30% to any other genomic sequence in the same genome.
- 11. (Currently amended) The method according to claim 1, wherein said amplified sequence-the printed product contains over 90 percent correct predetermined sequence.
- 12. (Previously presented) The method according to claim 1, wherein said array has a rectilinear format.

13-26. (Canceled)

27. (Previously presented) The method according to claim 1, wherein said predetermined gDNA sequence within the 3'UTR has a length of up to about 2000 nucleotides.

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28. (Currently amended) A method for amplifying expressed genetic sequences from genomic DNA (gDNA) selected from a higher-order eukaryotic species, for printing on DNA microarrays, wherein the method comprises:

identifying an exon of a gene defined by computer software;

selecting a predetermined gDNA sequence within the exon;

designing a probe for said predetermined gDNA sequence;

performing a first polymerase chain reaction (PCR) for the exon on gDNA to generate a first PCR-product;

separating the resultant-first PCR-product by a size-differentiation process selected from the group consisting of electrophoresis and chromatography;

selecting a predetermined band from the size-differentiated samples; performing a second PCR to amplify a product in the predetermined band; and depositing a sequence amplified by said second PCR to a substrate of an array printing the product of the second PCR on a substrate to form an array.

29. (Withdrawn) A method for making a DNA array, comprising:

performing a first PCR to amplify a 3'UTR, or a segment thereof, in a gDNA of a higher-order eukaryotic species;

separating products of said first PCR to select a product with a predetermined size;

performing a second PCR to amplify a sequence in said selected product; and depositing said amplified sequence to a substrate of the DNA array.

30. (Withdrawn) The method of claim 29, comprising:

performing PCRs to amply a plurality of 3'UTRs, or segments thereof, in genomic DNAs of said higher-order eukaryotic species;

separating products of said PCRs to select products with predetermined sizes; performing PCRs to amplify sequences in said selected products; and depositing said amplified sequences to the DNA array.

31. (Withdrawn) The method of claim 30, wherein each said 3'UTR is located between a stop codon and a polyadenylation signal of a different respective gene.

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- 32. (Withdrawn) The method of claim 31, wherein each said 3'UTR or segment comprises from about 75 to about 2,000 nucleotides, and each said separating step is accomplished by electrophoresis or chromatography.
- 33. (Withdrawn) The method of claim 31, wherein said higher-order eukaryotic species is a mammal, and each said 3'UTR or segment has an overall homology of no more than about 40% to any other genomic sequence in the genome of said mammal.
- 34. (Withdrawn) The method of claim 29, wherein said first and second PCRs are performed using the same pair of primers.